WEST Search History

Hide Items Restore Clear Cancel

DATE: Friday, December 14, 2007

Hide? Set Name Query

Hit Count

 $DB = PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD; \ PLUR = YES; \ OP = ADJ$

[i]

CB/ior-CEA.1

4

END OF SEARCH HISTORY

ClinicalTrials.gov PubMed Central

A service of the <u>U.S. National Lib</u> and the <u>National Ins</u>	My NEBlicine ?
--	----------------

All Databases I Journals Books	PubMed Nucleo	tide Protein	Genome	Structure	OMIM PMC	
Search PubMed	for				Preview	
ADOUT CITTLEZ Text Version	Limits Prev	riew/Index F	listory Clip	oboard Deta	ills	
Entrez PubMed Overview Help FAQ Tutorials New/Noteworthy E-Utilities	Search nTo save	umbers may n search indefin bine searches u	ot be continuitely, click qu	uery # and sele	activity. hes are represented ect Save in My NC #3 or click query #	CBI.
	Search	Mos	st Recent Qu	ieries	Time	Result
PubMed Services	#2 Search	Gailondo			12:25:24	<u>0</u>
Journals Database MeSH Database Single Citation Matcher Batch Citation Matcher	#1 Search	CB/ior-CEA	.1		12:24:50	0
Clinical Queries Special Queries LinkOut My NCBI	Clear History					
Related Resources Order Documents NLM Mobile NLM Catalog					·	
TOXNET Consumer Health Clinical Alerts						

Write to the Help Desk

NCBI | NLM | NIH

Department of Health & Human Services

Privacy Statement | Freedom of Information Act | Disclaimer

A service of the <u>U.S. National Librumy Ne slicine</u> and the <u>National Ins</u>	
--	--

All Databases I Journals Books	PubMed Nucleoti	de Protein	Genome	Structure	OMIM	PMC	
Search PubMed Go Clear	for	CB/ior-CEA.1				Preview	
Text Version	Limits Previe	ew/Index His	tory Clip	board De	tails_		•
Entrez PubMed Overview Help FAQ Tutorials New/Noteworthy E-Utilities	Search numTo save search	story will be lombers may not earch indefinite ne searches use s.	t be continu ely, click qu	ous; all sear ery # and se	ches are re lect Save	in My N	CBI.
	Search	Most	Recent Qu	eries		Time	Result
PubMed Services Journals Database MeSH Database Single Citation Matcher Batch Citation Matcher Clinical Queries Special Queries LinkOut My NCBI	#1 Search (CB/ior-CEA.1				12:24:50	<u>0</u>
Related Resources Order Documents NLM Mobile NLM Catalog NLM Gateway TOXNET Consumer Health Clinical Alerts ClinicalTrials.gov PubMed Central							
. asimod contrar			•				

Write to the Help Desk

NCBI | NLM | NIH

Department of Health & Human Services

Privacy Statement | Freedom of Information Act | Disclaimer

Welcome to STN International! Enter x:x LOGINID: SSPTALAB1643 PASSWORD: TERMINAL (ENTER 1, 2, 3, OR ?):2 Welcome to STN International Web Page for STN Seminar Schedule - N. America NEWS LMEDLINE coverage updated NEWS JUL 02 NEWS JUL 02 SCISEARCH enhanced with complete author names NEWS JUL 02 CHEMCATS accession numbers revised CA/CAplus enhanced with utility model patents from China NEWS 5 JUL 02 CAplus enhanced with French and German abstracts NEWS JUL 16 NEWS JUL 18 CA/CAplus patent coverage enhanced USPATFULL/USPAT2 enhanced with IPC reclassification NEWS JUL 26 NEWS 9 JUL 30 USGENE now available on STN AUG 06 CAS REGISTRY enhanced with new experimental property tags NEWS 10 NEWS 11 AUG 06 FSTA enhanced with new thesaurus edition NEWS 12 AUG 13 CA/CAplus enhanced with additional kind codes for granted patents CA/CAplus enhanced with CAS indexing in pre-1907 records NEWS 13 AUG 20 NEWS 14 AUG 27 Full-text patent databases enhanced with predefined patent family display formats from INPADOCDB NEWS 15 AUG 27 USPATOLD now available on STN NEWS 16 AUG 28 CAS REGISTRY enhanced with additional experimental spectral property data STN AnaVist, Version 2.0, now available with Derwent NEWS 17 SEP 07 World Patents Index NEWS 18 SEP 13 FORIS renamed to SOFIS INPADOCDB enhanced with monthly SDI frequency NEWS 19 SEP 13 CA/CAplus enhanced with printed CA page images from NEWS 20 SEP 17 1967-1998 CAplus coverage extended to include traditional medicine NEWS 21 SEP 17 patents EMBASE, EMBAL, and LEMBASE reloaded with enhancements SEP 24 NEWS 22 CA/CAplus enhanced with pre-1907 records from Chemisches NEWS 23 OCT 02 Zentralblatt NEWS 24 OCT 19 BEILSTEIN updated with new compounds NEWS 25 NOV 15 Derwent Indian patent publication number format enhanced NEWS 26 NOV 19 WPIX enhanced with XML display format NOV 30 ICSD reloaded with enhancements NEWS 27 NEWS 28 DEC 04 LINPADOCDB now available on STN 19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2, NEWS EXPRESS CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007. STN Operating Hours Plus Help Desk Availability NEWS HOURS

Enter NEWS followed by the item number or name to see news on that specific topic.

Welcome Banner and News Items

NEWS LOGIN

NEWS IPC8

For general information regarding STN implementation of IPC 8

STN maintenance downtime to be extended

The normal maintenance downtime for STN will be extended on December 15. STN will be unavailable beginning Saturday, December 15, at 17:00 U.S. Eastern Standard Time until Sunday, December 16, at 01:00.

The normal schedule for STN maintenance downtime (22:00 to 01:00) will resume on December 22.

All use of STN is subject to the provisions of the STN Customer agreement. Please note that this agreement limits use to scientific research. Use for software development or design or implementation of commercial gateways or other similar uses is prohibited and may result in loss of user privileges and other penalties.

FILE 'HOME' ENTERED AT 12:31:24 ON 14 DEC 2007

=> file caplus COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'CAPLUS' ENTERED AT 12:31:39 ON 14 DEC 2007 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2007 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 14 Dec 2007 VOL 147 ISS 26 FILE LAST UPDATED: 13 Dec 2007 (20071213/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> s CB/ior-CEA.1 'IOR-CEA.1' IS NOT A VALID FIELD CODE L1 0 CB/IOR-CEA.1

=> s (CB/ior CEA.1)
MISSING OPERATOR

```
23 CEA.1
                  (CEA(W)1)
             0 (CB/IOR AND CEA.1)
L2
=> s (CB and ior and CEA.1)
         14051 CB
          4649 CBS
         18336 CB
                  (CB OR CBS)
           192 IOR
             7 IORS
           198 IOR
                  (IOR OR IORS)
          6890 CEA
           204 CEAS
           7071 CEA
                  (CEA OR CEAS)
       9418668 1
             23 CEA.1
                  (CEA(W)1)
L3
              3 (CB AND IOR AND CEA.1)
=> duplicate remove L3
PROCESSING COMPLETED FOR L3
               3 DUPLICATE REMOVE L3 (0 DUPLICATES REMOVED)
=> d L4 bib abs 1-3
     ANSWER 1 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN
L4
     2003:892815 CAPLUS
AN
DN
     139:380012
TI
     Specific antibody fragments for the human carcinoembryonic antigen (CEA)
     Gavilondo Cowley, Jorge Victor; Ayala Avila, Marta; Freyre Almeida, Freya
IN
     de los Milagros; Acevedo Castro, Boris Ernesto; Bell Garcia, Hanssel;
     Roque Navarro, Lourdes Tatiana; Gonzalez Lopez, Luis Javier; Cremata
     Alvarez, Jose Alberto; Montesino Segui, Raquel
     Centro de Ingenieria Genetica y Biotecnologia, Cuba
PA
SO'
     PCT Int. Appl., 49 pp.
     CODEN: PIXXD2
DT
     Patent
LΑ
     Spanish
FAN.CNT 1
     PATENT NO.
                          KIND
                                  DATE
                                               APPLICATION NO.
                                               ______
                                                                        _____
     -----
                          _ _ _ _
                                  -----
                           A2
                                  20031113
                                               WO 2003-CU5
                                                                        20030428
ΡI
     WO 2003093315
     WO 2003093315
                          A3
                                  20040108
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
             PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT,
             TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                             CA 2003-2482411
                           A1
                                  20031113
                                                                        20030428
     CA 2482411
                                               AU 2003-223831
     AU 2003223831
                           A1
                                  20031117
                                                                        20030428
                                               BR 2003-4649
     BR 2003004649
                           A
                                  20040720
                                                                        20030428
     EP 1505076
                           A2
                                  20050209
                                              EP 2003-720119
                                                                        20030428
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                            CN 2003-809658
                                                                        20030428
     CN 1649901
                           Α
                                  20050803
                           Т
                                              JP 2004-501454
                                                                        20030428
     JP 2006500913
                                  20060112
```

	RU	2294939	C2	20070310	RU	2004-134601	20030428
	ZA	2004008720	A ´	20060329	ZA	2004-8720	20041027
	MX	2004PA10695	A	20050217	MX	2004-PA10695	20041028
	US	2005158322	A1	20050721	US	2005-511794	20050317
	US	2007199078	A1	20070823	US	2007-731442	20070330
PRAI	CU	2002-86	A	20020429			
	WO	2003-CU5	W	20030428			
	US	2005-511794	A3	20050317			•

AB The invention relates to mono- and bivalent (diabody) single-chain Fv-type (scFv) antibody fragments which are obtained using recombinant DNA techniques from the carcinoembryonic anti-antigen (CEA) monoclonal antibody (McA) CB/ior-CEA.1. The aforementioned McA has a high affinity for the CEA and is used in the diagnosis and monitoring of colorectal tumors in humans. As with the original McA, diabody and monovalent scFv fragments exhibit high affinities for the human CEA and recognize an epitope that is dependent on carbohydrate conservation. The diabody and monovalent scFv fragments have affinity consts. for the CEA of $(5.0 \pm 0.4) \times 109$ L mol-1 and $(2.8 \pm 0.4) \times 109$ L mol-1 a $0.3) \times 1010 \text{ L mol-1}$ resp. The two aforementioned fragments do not display cross-reactivity with normal human tissues and cells, except for the normal colonic mucosa where the CEA is occasionally present. fragments can be produced through expression in recombinant micro-organisms from the cloning of nucleic acid sequences that code for variable regions obtained from the hybridoma that is produced by the CB/ior-CEA.1 McA. As with the original McA, the diabody and the monovalent scFv have a capacity for the in vivo identification in rats of human CEA-producing cells which grow forming tumors. The monovalent scFv and diabody do not possess Fc domains and the mol. sizes of said monovalent scFv and diabody are 5 and 2.5 times, resp., less than the rat McA. As a result, the aforementioned monovalent scFv and diabody can better penetrate tissues in vivo and are less immunogenic in humans.

- L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 1992:122152 CAPLUS
- DN 116:122152
- TI Primer design for the cloning of immunoglobulin heavy-chain leader-variable regions from mouse hybridoma cells using the PCR
- AU Coloma, Maria J.; Larrick, James W.
- CS Genelabs, Inc., Redwood City, CA, 94063, USA
- SO BioTechniques (1991), 11(2), 152-4, 156 CODEN: BTNQDO; ISSN: 0736-6205
- DT Journal
- LA English
- AB To facilitate the rapid cloning and sequencing of rearranged murine heavy-chain variable regions, a set of universal primers was designed using conserved sequences of leader (signal peptide), framework one and constant regions of the Ig heavy-chain genes. RNA was extracted from the mouse hybridoma cells secreting monoclonal antibodies: IOR-T3 (anti-CD3), C6 (anti-P1 of N. meningitidis B385), IOR-T1 (anti-CD6), CB-CEA.1 (anti-carcinoembryonic antigen), CB-Fib.1 (anti-human fibrin) and CB-Hep.2 (anti-hepatitis B surface antigen). First-strand cDNA was synthesized and amplified using PCR. The primers successfully amplified correct size fragments from cDNA prepared from all hybridomas. These methods will facilitate the cloning and sequencing of mouse Ig variable regions.
- L4 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 1991:56891 CAPLUS
- DN 114:56891
- TI Specific amplification of rearranged immunoglobulin variable region genes from mouse hybridoma cells
- AU Gavilondo-Cowley, Jorge V.; Coloma, Maria J.; Vazquez, Javier; Ayala, Marta; Macias, Amparo; Fry, Kirk E.; Larrick, James W.

CS Div. Hybridomas Anim. Models, Cent. Genet. Eng. Biotechnol., Havana, Cuba

Hybridoma (1990), 9(5), 407-17 CODEN: HYBRDY; ISSN: 0272-457X

DT Journal

SO

LΑ English

AB This article describes how the polymerase chain reaction (PCR) and primers designed for conserved sequences of leader (L), framework one (FR1) and constant (CONST) regions of Ig light and heavy chain genes can be used for the cloning and sequencing of rearranged antibody variable regions from mouse hybridoma cells. RNA was extracted from the mouse hybridoma cells secreting MAbs: IOR-T3a (anti-CD3), C6 (anti-P1 of Neisseria meningitidis B385), IOR-T1 (anti-CD6), CB-CEA .1 (anti-carcinoembryonic antigen), and CB-Fib.1 (anti-human fibrin). First strand cDNA was synthesized and amplified using PCR. The newly designed primers are superior to others reported recently in the literature. Isolated PCR DNA fragments of C6 and IOR-T3a were sequenced after asym. amplification, or M13 cloning. The FR1/CONST primer combinations selectively amplified mouse light chains of groups kappa II, V, and VI, and heavy chains of groups IIa and IIc. The L/CONST primers for light chains amplified light chains from all 4 hybridomas. The methods greatly facilitate structural and functional studies of antibodies by reducing the efforts required to clone and sequence their variable regions.